

Blood Tolerant Laccase by Directed Evolution

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SUMMARY

High-redox potential laccases are powerful biocatalysts with a wide range of applications in biotechnology. We have converted a thermostable laccase from a white-rot fungus into a blood tolerant laccase. Adapting the fitness of this laccase to the specific composition of human blood (above neutral pH, high chloride concentration) required several generations of directed evolution in a surrogate complex blood medium. Our evolved laccase was tested in both human plasma and blood, displaying catalytic activity while retaining a high redox potential at the T1 copper site. Mutations introduced in the second coordination sphere of the T1 site shifted the pH activity profile and drastically reduced the inhibitory effect of chloride. This proof of concept that laccases can be adapted to function in extreme conditions opens an array of opportunities for implantable nanobiodevices, chemical syntheses, and detoxification.

INTRODUCTION

The extension of an enzymes' capacity to function in extreme environmental conditions is an issue that promises to deliver countless benefits. Due to their extraordinary versatility, the study of the ligninolytic enzymatic consortium of oxidoreductases secreted by white-rot fungi is of special biotechnological interest—mainly high-redox potential laccases (HRPLs), peroxidases, and H₂O₂-supplying enzymes (Martínez et al., 2009). However, the absence of catalytic activity at neutral/basic pHs, along with the inhibition by modest concentrations of different substances (halides, metal ions, fatty acids, detergents), remain a serious obstacle to their further exploitation. HRPLs (EC 1.10.3.2) that are active under such inclement conditions would be very desirable to be used in applications that range from organic synthesis to bioremediation (Alcalde, 2007; Gianfreda et al., 1999). In addition, HRPLs belong to the exclusive group of oxidoreductases that are capable of accepting electrons directly from the cathode of a biofuel cell or an amperometric biosensor. Indeed, the set of advantages that HRPLs have to offer (i.e., high current densities, direct electron transfer, low

overpotential for O₂ reduction, and high operational stability) situate them among the best suited candidates for enzyme-based bioelectronic devices (Shleev and Ruzgas, 2008). Possibly, one of the most attractive challenges in this field focuses on achieving implantable self-contained wireless 3D-nanobiodevices that work in different physiological fluids (blood, saliva, tears). Such nanobioelectronic devices are comprised of a biosensor array to detect different metabolites in vivo (glucose, insulin), a transmitter/transducer to externalize the information and a biofuel cell to power the entire system. The main shortcomings in the engineering of this type of technology stem from the difficulties in miniaturizing their individual elements (antenna, transducer), and in tailoring reliable and stable enzymes to catalyze the biocathode reaction in which the O₂ dissolved in the fluids is reduced to H₂O (Bullen et al., 2006; Castillo et al., 2004; Kim et al., 2006). Unfortunately, HRPLs are inactive at blood pH (~7.4) and they are also strongly inhibited by chloride concentrations much lower than those found in blood (140–150 mM), which limits this specific application and the use in other areas where basic pHs or chloride ions are presented (i.e., dye-stuff processing, waste-water treatment, pollutant remediation, pharmaceutical compound synthesis, and food processing, to name a few) (Alcalde, 2007; Gianfreda et al., 1999; Riva, 2006; Rodgers et al., 2010).

Here, we report the engineering by directed evolution of an HRPL that is functional in human blood. The ultimate mutant enzyme obtained through this evolutionary process was characterized comprehensively and its evolved features were tested on real human blood samples, revealing the mechanisms underlying this relevant improvement.

RESULTS AND DISCUSSION

Laboratory Evolution Approach

Our point of departure in this study was a HRPL (OB-1 mutant) previously tailored in our laboratory by in vitro evolution and semirational approaches to be active, soluble, highly thermostable, and readily secreted by yeast (Maté et al., 2010). To convert OB-1 to an active blood-tolerant laccase, we designed a screening assay based on human blood composition. As real plasma or blood are unsuitable for screening mutant libraries in the experimental context of directed evolution, we developed a surrogate complex medium ("blood-buffer") mimicking the biochemical composition of human blood. This buffer contained a laccase colorimetric substrate but it lacked any cells and coagulating agents. Three consecutive rescreens were conducted to

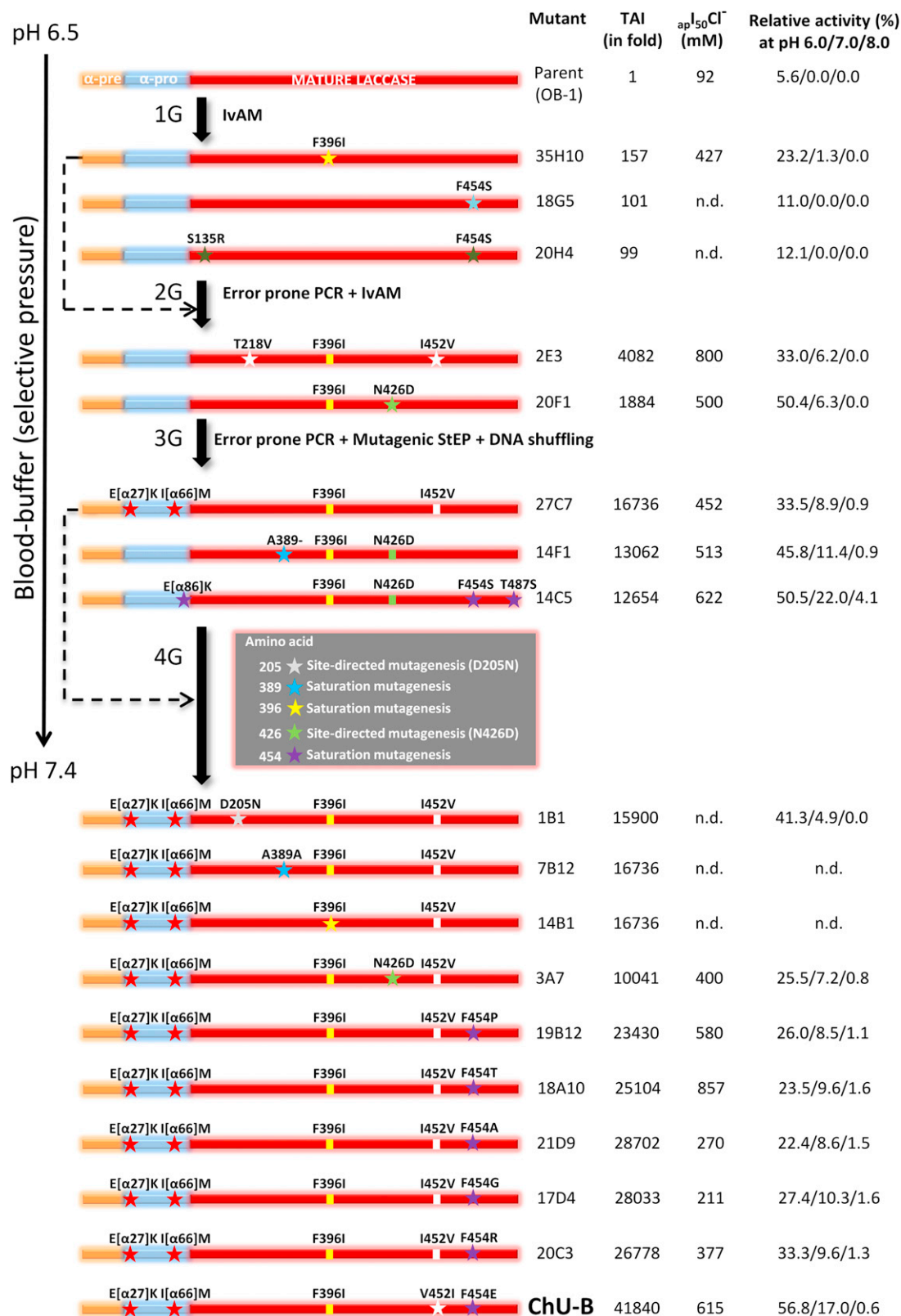


Figure 1. Directed Laccase Evolution

The parental type fusion gene (OB-1 mutant) (Maté et al., 2010) is formed by the α -factor prepro-leader, replacing the original laccase signal sequence to improve secretion in *S. cerevisiae*, and the mature laccase. The α -factor preleader is represented in orange, the α -factor proleader in blue and the mature laccase in red. (legend continued on next page)

rule out the selection of false positives. Like all HRPLs, the OB-1 mutant exhibits neither activity nor significant internal electron transfer at pH 7.4 (Maté et al., 2010). Accordingly, for the first generation the pH of the blood-buffer was set at 6.5, resulting in a decrease in laccase activity of over 90%. In successive generations selective pressure was progressively enhanced until physiological pH values were reached. We took advantage of the high frequency of homologous DNA recombination of the *Saccharomyces cerevisiae* machinery to create diversity (Alcalde, 2010; Gonzalez-Perez et al., 2012). Thus, in each evolution cycle different DNA-recombination methods (both in vivo and in vitro) were combined to enhance the complexity of the mutant libraries in this pool. The last round of evolution was dedicated to the detailed rational evaluation of several positions that produced substantial improvements in the total activity by saturation mutagenesis, site-directed mutagenesis, and mutational recovery (Figure 1; Table S1 and Supplemental Results available online). When the last variant of the evolutionary pathway was obtained (ChU-B mutant), it exhibited an increase in total activity in blood-buffer over 40,000-fold versus the parental type.

Laccase Mutant in Blood

The behavior of the mutant laccase was tested in human plasma and blood. Oxygen consumption in physiological fluids enriched with ascorbic acid (a poor laccase substrate naturally present in blood) was monitored with a Clark electrode, revealing comparable responses for plasma and blood (185 and 127 min⁻¹, respectively; Figure 2A). The 1.5-fold increase in ChU-B activity in plasma suggested that blood cells interfere with the detection method. To confirm that the enzyme was active under physiological conditions regardless of the reducing substrate used, oxygen reduction against common laccase substrates (2,2'-azino-bis(3-ethylbenzothiazoline-6-sulphonic acid) (ABTS), K₄[Fe(CN)₆]) was measured in blood-buffer, obtaining similar values for all compounds tested (Figure 2A).

Laccases contain four catalytic copper (Cu) ions arranged at two different sites: the T1 site, at which the substrate is oxidized, and a trinuclear Cu cluster (with one Cu T2 and two Cu T3), at which oxygen is reduced to water (Morozova et al., 2007). If the laccase is properly connected to an electrode via the T1 Cu, the electroactive surface of the device can replace the electron-donating natural substrates of the enzyme. Redox titration of the T1 Cu site was performed at physiological pH using a microspectroelectrochemical cell with a gold capillary electrode (Figure 2B). The ChU-B mutant exhibited a high redox potential (+720 mV versus NHE) at pH 7.4, and it reversibly cycled between its fully oxidized and fully reduced state. To further verify that the mutant was electrochemically active under physiological conditions, ChU-B was adsorbed onto low-density graphite electrodes

and the electrocatalytic response of O₂ reduction by direct electron transfer was recorded at pH 7.4 (Figure 2C). No responses were detected for any of the HRPLs tested at this pH.

Inhibition by Halides and Hydroxides

The novel properties of ChU-B were dissected in terms of its activity versus pH and halide inhibition. The pH activity profiles against phenolic (2,6-dimethoxyphenol) (DMP) and nonphenolic (ABTS) compounds revealed a notable shift toward less acidic values, including a change in DMP optimum activity pH from 4.0 to 5.0–6.0 (Figures 3A, 3B, S1A, and S1B). At pH 7.0, ChU-B retained ~50% and ~20% of its activity for DMP and ABTS, respectively, whereas the parental type displayed negligible activity under these conditions. Similarly, ChU-B activity at pH 6.0 was over 90% and 50% for DMP and ABTS, respectively, whereas that of the parental type was ~20%. This drastic improvement in activity at near neutral pH led to a small but noticeable increase in ChU-B activity even at pH 8.0. To date, the engineering of chimeric laccases had only succeeded in slightly shifting the pH profile, at the expense of sacrificing redox potential (Cusano et al., 2009). By contrast, ChU-B variant exhibits activity at neutral/alkaline pH values while keeping a high redox potential at the T1 Cu site.

It is well known that increases in pH inhibit laccase activity, due to a decrease in H⁺ availability and the binding of OH⁻ ions to the T2 Cu, which in our laccase is tricoordinated with His64, His397, and one H₂O molecule (Matera et al., 2008; Xu, 1997). HRPLs are also inhibited by halides (fluoride, chloride, and bromide, but not iodide), with an inhibitory potency inversely proportional to the diameter of the anion (F⁻ > Cl⁻ > Br⁻). This dependence is possibly due to an access limitation for the binding of the larger halide ions to the laccase catalytic sites (Xu, 1996, 1997). The I₅₀ measured for the different halides (the concentration of halide at which the enzyme retains 50% of its initial activity) was independent of the substrate employed (Table S2). The I₅₀ of ChU-B for Cl⁻ improved from 176 to 1,025 mM with ABTS as substrate (Figure 3C), to the best of our knowledge the highest I_{50Cl⁻} reported for any basidiomycete HRPL. It was not possible to measure Br⁻ inhibition for ChU-B, as the I₅₀ of the parental type was already above 1,300 mM, impeding measurement by surpassing the limit of substrate solubility at such high salt-concentrations. As expected, there was strong inhibition in the presence of F⁻ (in the μM range), and although F⁻ inhibition was not a goal of the directed evolution study, the I_{50F⁻} was slightly enhanced (from 70 to 109 μM, Figure 3D). Halide inhibition was also measured at physiological pH values and ChU-B was not sensitive to increasing concentrations of chloride (ranging from 100 to 800 mM) (Figure 3E). At pH 7.4, very little H₂O dissociation was observed, although the resulting 251 nM of OH⁻ may be enough to

New mutations are depicted as stars and accumulated mutations as squares; A389- is a deleted residue. TAI (total activity improvement in blood-buffer): a value indicating the improvement in laccase activity detected in *S. cerevisiae* microcultures for each mutant compared with the parental OB-1. Measurements were performed in quintuplicate from supernatants of independent cultures grown in 96-well plates, using 3 mM ABTS as substrate. The *ap*I_{50Cl⁻} (apparent I₅₀ for NaCl) indicates the concentration of NaCl at which the enzyme retains 50% of its ABTS-activity, as determined in *S. cerevisiae* microcultures for each mutant after a 2 hr incubation. Relative activity at different pH values was assessed with 3 mM ABTS as substrate. Dashed arrows indicate the mutant used as a parent in each generation. N.d., not determined. Silent mutations are not included. Over 10,000 clones were screened in four rounds of molecular evolution combining in vivo (IvAM, DNA shuffling, IVOE) and in vitro (error-prone PCR, mutagenic STEP) recombination/mutagenic methods. In the last cycle of evolution, the final variant (ChU-B) was obtained after analyzing several positions by site-directed mutagenesis and saturation mutagenesis.

See also Supplemental Results, Tables S1 and S5, and Figure S3.

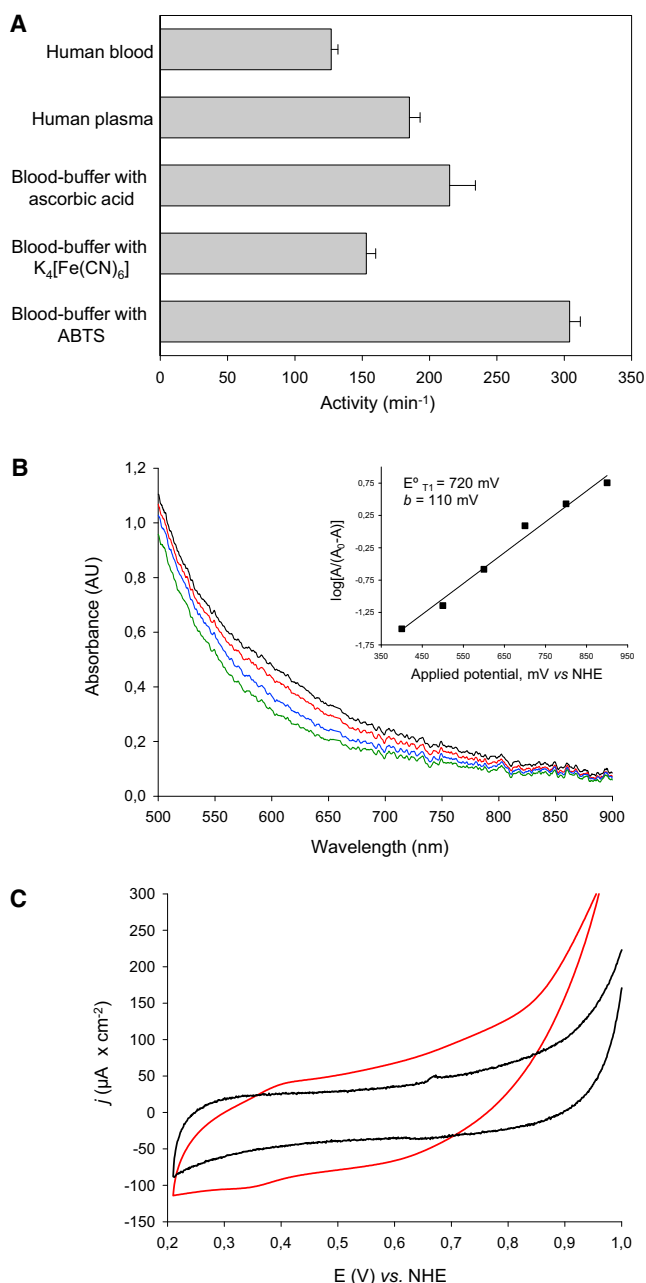


Figure 2. Activity in Physiological Fluids and Spectro-Electrochemical Characterization

(A) Activity of the ChU-B mutant determined by measuring O₂ consumption. To monitor laccase activity in blood and plasma, the fluids were supplemented with 10 mM ascorbic acid and the pH adjusted to 7.4 prior to adding the enzyme. Each bar, including the SD, is from three independent experiments. (B) Redox titration of ChU-B at physiological pH values. Each redox titration was carried out in both directions: from the fully oxidized to the fully reduced state of the enzyme (reductive titration) and vice versa (oxidative titration). The spectra of laccases were recorded at redox equilibrium and typical spectra of the oxidized, partly reduced and fully reduced enzymes are depicted. Equilibration of the T1 copper center at each potential applied was apparent from the stabilization of absorbance at 600 nm. As the redox mediators used are transparent above 500 nm, the spectral changes at 600 nm were attributed to the T1 Cu of the laccase. Spectra from the reductive titration, corresponding to oxidized laccase (black curve, applied potential +1,000 mV versus NHE),

efficiently bind at the T2 Cu site and promote the inhibition of native HRPLs. The gradual increase in blood-buffer pH throughout evolution reflects the [OH⁻] increase and [H⁺] decrease, and it permitted a selection of variants that are more tolerant to both hydroxyl and chloride ions in the same screening assay. Comparing ChU-B with other HRPLs revealed that, depending on the substrate, our variant was 12- to 20-fold less sensitive to F⁻ than the HRPLs from *Trametes trogii* and *Trametes villosa* (Garzillo et al., 1998; Xu, 1996). Similarly, the I_{50Cl⁻} of ChU-B was 26- to 164-fold higher than corresponding HRPLs. The ascomycete laccase from *Botrytis aclada* exhibited the highest I_{50Cl⁻} described to date (1.4 M with DMP as substrate) (Kittl et al., 2012), yet its negligible activity at neutral pH and poor stability precludes its use in implantable biocathodes or other environmental processes. Comparing ChU-B with laccases from bacterial sources revealed a 1.5-fold better chloride tolerance than that of the halide-resistant polyphenol oxidase from *Marinomonas mediterranea* (Jimenez-Juarez et al., 2005). This halide tolerance of the ChU-B mutant is remarkable given that, as a rule, fungal laccases are much more sensitive to halide inhibition than low-redox potential laccases from bacteria (Niladevi et al., 2008; Singh et al., 2011).

We assessed maximum turnover rates of ChU-B in blood-buffer. The synergic effect that improved activity at physiological pH in association with a strong chloride tolerance resulted in initial rates of 427 and 143 mol substrate/min/mol enzyme for DMP and ABTS, respectively. In equivalent conditions, the parental type showed almost no activity. Significantly, the turnover rates of the ChU-B mutant was 3- to 4-fold lower than that of the parental type under optimal conditions (i.e., acidic pH and in the absence of inhibitors; Table S3). Intrigued by these results, we analyzed mutations in the laccase structure.

Structural Analysis of Mutations

The laccase catalytic scaffold is formed by the T1 Cu site close to the surface, although the copper is not solvent-exposed. The T1 Cu is trigonally coordinated by His455, His394, and Cys450, the latter forming part of the highly-conserved His451-Cys450-His449 tripeptide, which connects to the trinuclear cluster located 12 Å away. The electrons from T1 Cu are transferred through two intermolecular electron transfer pathways governed by the aforementioned tripeptide to the T2/T3 trinuclear Cu cluster, where O₂ binds and is reduced to two molecules of H₂O. This second active site is buried deep in the laccase structure, with a T2 Cu

partly reduced enzymes (red and blue curves, +700 mV and +600 mV versus NHE, respectively) and the fully reduced laccase (green curve, +300 mV versus NHE) are shown. Inset: a typical Nernst plot of the dependence of the applied potential versus absorbance at 600 nm, and averaged parameters calculated from the reductive and oxidative titrations (midpoint value of +720 mV versus NHE and a slope of 110 mV).

(C) Cyclic voltammograms of O₂ reduction obtained using a polished, low-density graphite (LDG) electrode (black), and a LDG electrode with ChU-B laccase adsorbed to its surface (red). Measurements were carried out in a three-electrode electrochemical cell filled with 100 mM sodium phosphate buffer pH 7.4 and after bubbling O₂ at a pressure of 1 atm for 15 min, using a platinum wire as the counter electrode and a BAS Ag/AgCl reference electrode. Measurements were performed using Autolab PGSTAT30 controlled by GPES 4.9 software.

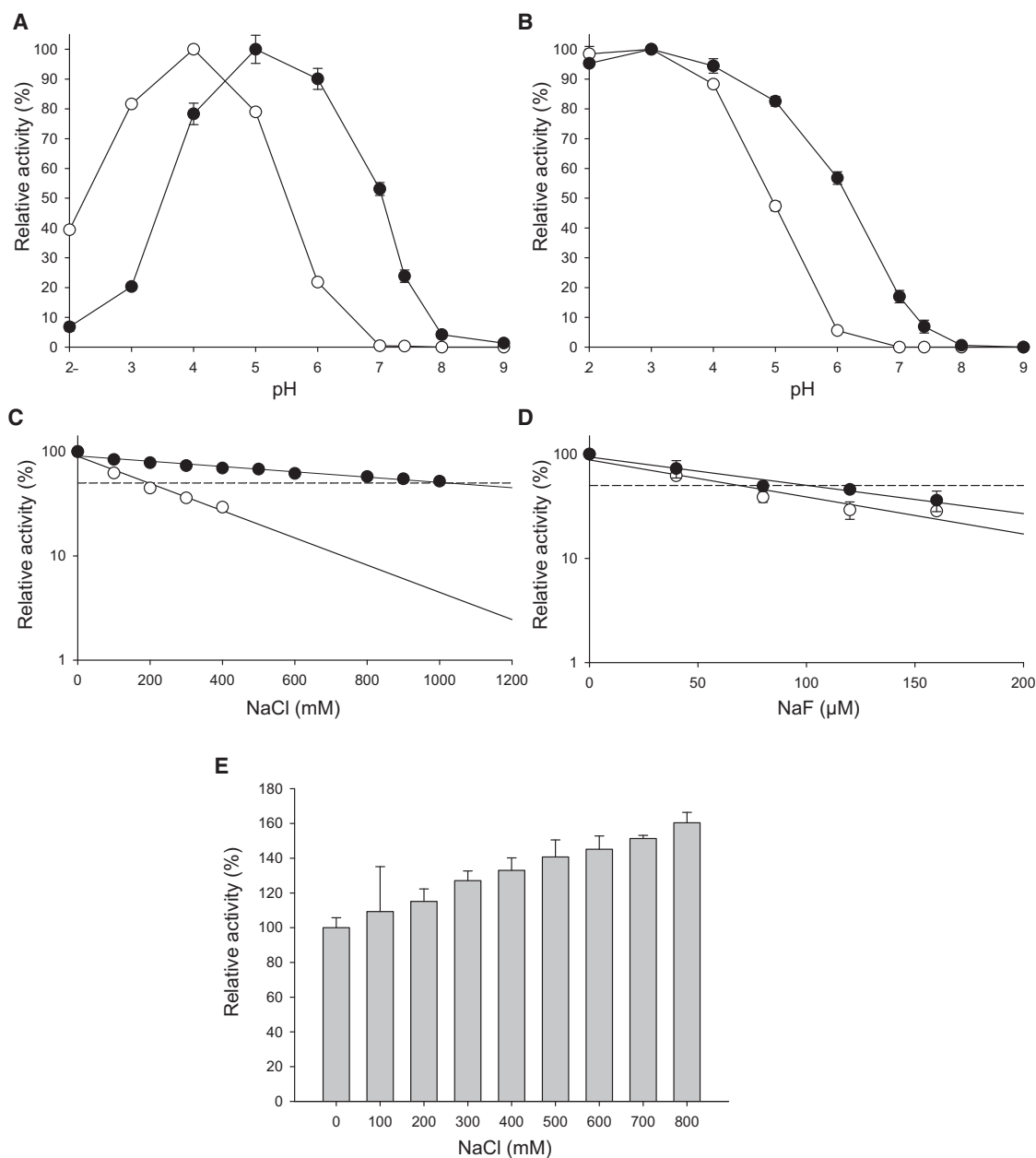


Figure 3. Biochemical Characterization

(A and B) pH activities profiles were measured in 100 mM Britton and Robinson buffer at different pH values with 3 mM DMP (A) or ABTS (B) as substrates. White circles, parental type; black circles, ChU-B mutant. Laccase activity was normalized to the optimum activity value.

(C–E) Inhibition by halides. (C) Cl⁻ inhibition measured with 2.4 mM ABTS in 100 mM sodium acetate buffer pH 4.0. (D) F⁻ inhibition measured with 2.4 mM ABTS in 100 mM sodium acetate buffer pH 4.0. (E) Cl⁻ inhibition of ChU-B measured with 2.4 mM ABTS in 100 mM sodium phosphate buffer pH 7.4. Each value represents the mean and SD derived from three independent experiments.

See also Figure S1 and Tables S2 and S3.

tricoordinated by two His residues, one molecule of water and two tetracoordinated T3 coppers. The entrance of O₂ and the exit of H₂O to the T2/T3 site occur via two solvent channels. Bearing in mind that halide/OH⁻ inhibition allegedly occurs at the T2/T3 cluster, interrupting the internal electron transfer (Naki and Varfolomeev, 1981; Naqui and Varfolomeev, 1980), we expected to find mutations located in the surroundings of

the trinuclear copper cluster, or in either of the two channels for oxygen and water transit (Matera et al., 2008; Xu et al., 1998). However, the two mutations introduced in the mature ChU-B (F396I and F454E) were located at the second coordination sphere of the T1 Cu site (Figure 4). Indeed, most of the mutations discovered during the course of evolution were placed in the same region (at an average distance of 7.5 Å

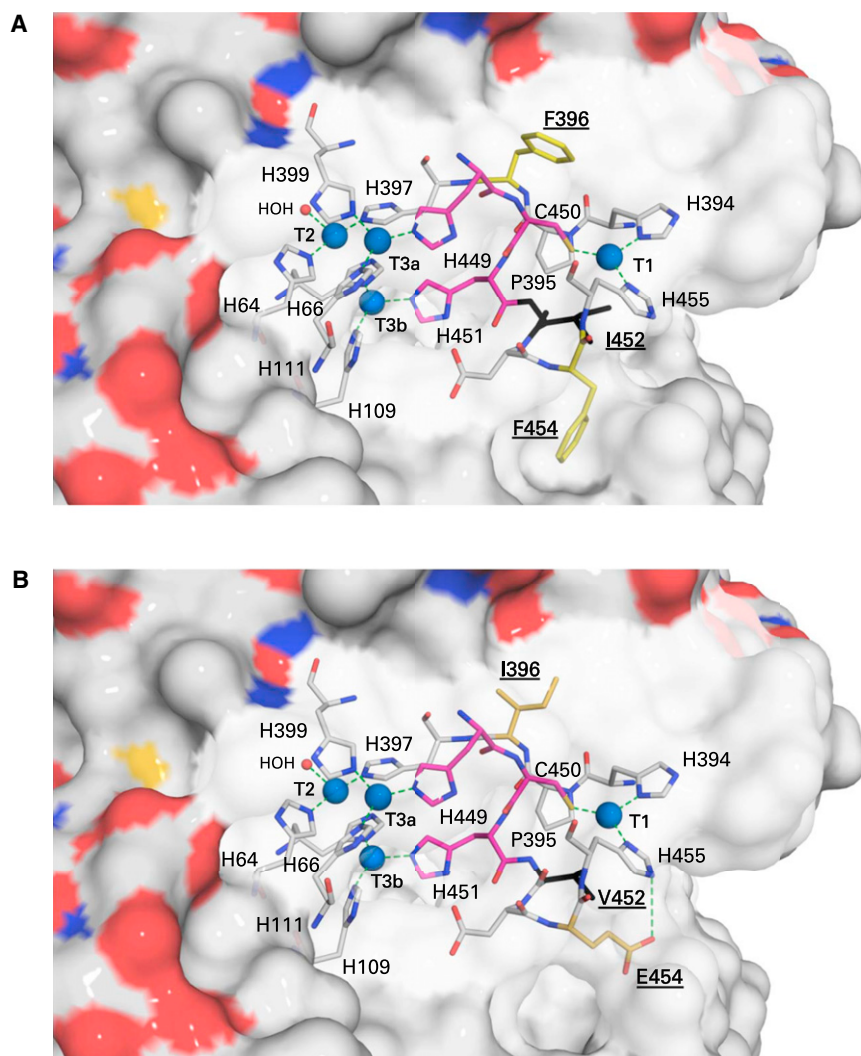


Figure 4. Mutations in Evolved Laccase

(A and B) Details of the mutations in the ChU-B variant (B) compared with the corresponding residues in the parental type (A). The F396I and F454E mutations are shown in yellow, and the I452V mutation (reverted in ChU-B) in black. Blue spheres represent copper atoms. The residues of the internal transfer pathway from T1 Cu to T2/T3 cluster are colored magenta. Residues involved in the first coordination sphere of the catalytic coppers and their interactions (as green dashes) are also represented. The electrostatic surface of the protein structure is shown in the background. The 3D structure model is based on the crystal structure of the *Trametes troglia* laccase (97% identity, PDB: 2HRG) (Matera et al., 2008). See also Figure S2 and Table S4.

to be established (Figure 4). It is generally accepted that electron transfer from the substrate to the T1 Cu is the rate-limiting step of laccase catalysis (Alcalde, 2007; Gianfreda et al., 1999). Modification of the second coordination sphere of the T1 Cu reduces activity at acidic pH values and simultaneously compensates for T2 Cu inhibition so that ChU-B maintains activity in the presence of halides and OH^- . These results are consistent with previous studies of an ascomycete low-redox potential laccase subjected to comprehensive site-directed mutagenesis at the T1 region (Xu et al., 1998). Recently, the parental type of the current research—the HRPL OB-1 mutant evolved for secretion and stability in yeast (Maté et al., 2010)—was analyzed by a computational algorithm to understand

from the T1 Cu site), indicating an important influence of this region on halide/ OH^- inhibition (Figure S2; Table S4).

F396I was discovered in the first generation, and it provoked the greatest improvement in activity of the entire evolutionary process (157-fold increase in activity over the parental type). Additional saturation mutagenesis at this position did not confer any further improvement (Figure 1). The highly conserved Phe396 plays a key role in the redox potential of the T1 site, and hence in HRPL catalysis (Matera et al., 2008). Phe396 acts as a bridge connecting the T1 Cu and T2/T3 Cu cluster via Pro395 (contiguous to the coordinating His394 of the T1 Cu) and His397, which coordinates the T2 Cu (Figure 4). The F454E mutation was generated by subjecting position 454 to saturation mutagenesis and subsequent screening, as some mutants discovered in the first and third generations already contained a mutation at this position (Figures 1 and S3). We selected six different mutants with F454P/T/A/G/R/E substitutions, which boosted activity in blood-buffer up to 2.5-fold (Supplemental Results). The F454E mutation is contiguous to the coordinating His455 of the T1 Cu. Analysis of our model revealed that upon mutation, an H-bond with the imidazole group of His455 appears

the physical forces that rule the stability of the variant (Christensen and Kepp, 2012). Indeed, the combination of *in silico* computational methods (based on Monte Carlo simulations and molecular dynamics) may provide a new twist in the study of this blood tolerant laccase. In particular, the use of quantum mechanics/molecular mechanics (QM/MM) methods—for mapping internal electron transfer pathways (Guallar and Wallrapp, 2008)—as well as the protein energy landscape exploration (PELE)—for assessing the traffic of halides and hydroxides through the laccase structure (Borrelli et al., 2005)—could be valuable approaches to discern the hidden physicochemical principles behind the laccase improvements.

SIGNIFICANCE

HRPLs are considered by many as the green biocatalysts of the 21st century: they easily oxidize hundreds of compounds using oxygen from air and release water as the only byproduct (Maté et al., 2011). Moreover, they can be immobilized onto electrodes and they mediate direct electron transfer, permitting their integration into nanobioelectronic devices.

Our blood-tolerant laccase displays all these properties, thereby enabling its incorporation into physiological fluids. We are currently designing a full wireless 3D-nanobiodevice prototype for use in human blood, containing an ad hoc transmitter and a biosensor powered by a biofuel cell, to which ChU-B is attached (S. S., M.A. A.D.L., R.L., M.P., M.F., D.M., D.G.P., W., Schuhmann, L. Gorton, D. Leech, P. Bartlett, M. Duarte, E. Magner, F. Sebelius, and D. Sutherland, unpublished data). This technology should permit less invasive and more reliable real-time in situ monitoring. Indeed, nanobiodevices can minimize the often-critical time to detect a potentially life threatening physiological imbalance and shorten the intervention time for medical teams. With an extraordinary resistance to halides and significant activity at neutral/alkaline pH values, the benefits of our evolved HRPL may also be extended to applications such as the development of medical bioassays, bioremediation (e.g., oxidation of pesticides, PAHs, contained-waste waters, dye processing), pulp-kraft biobleaching, organic syntheses, and cofactor/coenzyme regeneration, among other processes in which high pH and/or strong salt contents represent major hurdles (Alcalde et al., 2006; Rodríguez Couto and Toca Herrera, 2006; Kunamneni et al., 2008; Riva, 2006; Witayakran and Ragauskas, 2009).

EXPERIMENTAL PROCEDURES

All chemical reagents were of the highest purity commercially available. The oligonucleotides used along the evolutionary process (Table S5) were purchased from Isogen Life Science (De Meern, The Netherlands). Culture media were prepared as described previously (Maté et al., 2010).

Laboratory Evolution

For each generation, PCR fragments were cleaned, concentrated and loaded onto a low melting point preparative agarose gel (Bio-Rad, Hercules, CA), and then purified using the Zymoclean gel DNA recovery kit (Zymo Research, Orange, CA). PCR products were cloned under the control of GAL1 promoter of the pJRoC30 expression shuttle vector (kindly donated by Prof. F.H. Arnold from Caltech, CA), replacing the parent gene in pJRoC30. To remove the parent gene, the pJRoC30 plasmid was linearized with BamHI and XhoI (New England BioLabs, Hertfordshire, UK), and the linear plasmid was concentrated and amplified as described above for the PCR fragments.

First Generation: IvAM

Using the OB-1 mutant as the parental type (Maté et al., 2010), a library of over 3,000 clones was constructed by in vivo assembly of mutant libraries with different mutational spectra (IvAM) (Zumárraga et al., 2008). Independent mutagenic PCR reactions with Taq/MnCl₂ (Sigma-Aldrich, Madrid, Spain) and Mutazyme II DNA polymerase (Genemorph II Random mutagenesis kit, Stratagene, CA) were carried out on a gradient thermocycler (Mycycler, Bio-Rad) using the following parameters: 95°C for 2 min (1 cycle); 94°C for 0.75 min, 53°C for 0.75 min, 74°C for 3 min (28 cycles); 74°C for 10 min (1 cycle). In a final volume of 50 µl, the Taq/MnCl₂ PCR amplification contained 90 nM RMLN, 90 nM RMLC, 0.1 ng/µl template, 0.3 mM dNTPs (0.075 mM each), 3% DMSO, 1.5 mM MgCl₂, 0.01 mM MnCl₂, and 0.05 U/µl Taq polymerase. Mutazyme II amplification was carried out with 372 nM RMLN, 372 nM RMLC, 40 ng/µl template, 0.8 mM dNTPs (0.2 mM each), 3% DMSO, and 0.05 U/µl Mutazyme II per 50 µl of reaction product. Under these conditions the mutational rates of Taq/MnCl₂ and Mutazyme II libraries were 0–3 and 4.5–9 mutations per 1,000 bp, respectively. To promote in vivo ligation, overhangs of 40 and 66 bp homologous to the linear vector were designed. The two libraries were mixed in equimolar amounts and transformed into competent *S. cerevisiae* cells of the protease-deficient strain BJ5465 (LGC Promochem, Barcelona, Spain) using the yeast transformation kit (Sigma-Aldrich) together with the linearized vector (ratio vector/library 1:4).

Transformed cells were plated in synthetic complete drop-out plates and incubated for 3 days at 30°C. Colonies were then picked and subjected to the screening assay and to three successive rescreenings, as described below.

Second Generation: Error Prone PCR + IvAM

Using the 35H10 mutant from the first generation as the parental type, three mutant libraries (~700 clones each) were constructed and independently screened. The first library was prepared with Taq/MnCl₂, the second with Mutazyme II and the third with IvAM, as described for the first generation.

Third Generation: Mutagenic StEP and In Vivo DNA Shuffling

Two independent libraries were created and explored using the two best mutants from the second round, 2E3 and 20F1, as the parental types. For the first library, the parental types were independently prepared with Taq/MnCl₂ and the mutagenic products in vivo shuffled in *S. cerevisiae*. The second library was prepared by in vitro recombination through staggered extension process (StEP) (Zhao et al., 1998). The StEP-PCR reaction (50 µl final volume) contained 0.5 µM RMLN, 0.5 µM RMLC, 0.1 ng/µl of each DNA-template, 0.8 mM dNTPs (0.2 mM each), 3% DMSO, 1 mM MgCl₂, 0.01 mM MnCl₂, and 0.02 U/µl iProof high-fidelity DNA polymerase (Bio-Rad). The PCR conditions were as follows: 95°C for 2 min (1 cycle); 94°C for 0.5 min, 55°C for 0.33 min (90 cycles). An electrophoretic band of ~2 kb was purified and further subjected to mutagenic PCR with Mutazyme II. This mutagenic product was in vivo shuffled as described above.

Fourth Generation: Site-Directed and Saturation Mutagenesis Studies

Two individual mutants and three saturation mutagenesis libraries were constructed by in vivo overlap extension (IVOE) (Alcalde, 2010) using the 27C7 mutant as the parental template. The PCR reactions were carried out using the iProof high-fidelity DNA polymerase.

D205N Mutant. (Madzak et al., 2006) The primers for PCR 1 were RMLN and D205N-REV and for PCR 2 were D205N-FOR and RMLC.

N426D Mutant. The primers for PCR 1 were RMLN and N426D-REV and for PCR 2 were N426-FOR and RMLC.

Saturation Mutagenesis at Position 389. The primers for PCR 1 were RMLN and SAT389-REV and for PCR 2 were SAT389-FOR and RMLC.

Saturation Mutagenesis at Position 396. The primers for PCR 1 were RMLN and SAT396-REV and for PCR 2 were SAT396-FOR and RMLC.

Saturation Mutagenesis at Position 454. The primers for PCR 1 were RMLN and SAT454-REV and for PCR 2 were SAT454-FOR and RMLC.

Screening Assay and High-Throughput Protocol

Laccase mutant libraries were screened in a medium (blood-buffer) containing the colorimetric laccase substrate ABTS and that simulates the composition of human blood, although it lacks coagulating agents and cells. Blood-buffer composition: 150 mM NaCl, 18 mM NaHCO₃, 1 mg/ml D-(+)-glucose, 4.3 mM urea, 0.87 mM MgSO₄, 0.4 mM D-(+)-fructose, 0.1 mM L-cysteine, and 4.29 mM ABTS in 100 mM sodium phosphate buffer. The blood-buffer pH was set at 6.5 for the first generation and was gradually increased throughout evolution to reach physiological pH (7.4) in generation 4. The general HTP-screening protocol was as reported previously (Maté et al., 2010) with some modifications. Individual clones were picked and cultured in 96-well plates (Greiner Bio-One, Frickenhausen, Germany) containing 50 µl of minimal medium per well. In each plate, column number 6 was inoculated with the parental type and one well (H1-control) was left uninoculated. Plates were sealed to prevent evaporation and incubated at 30°C and 80% relative humidity on a shaker at 225 rpm (Minitron-INFORS, Biogen, Spain). After 48 hr, 160 µl of expression medium was added to each well and the plates were incubated for 24 hr. The plates (master plates) were then centrifuged (Eppendorf 5810R centrifuge, Germany) for 5 min at 3,000 × g at 4°C and 60 µl of the supernatant was transferred from the master plate onto a replica plate with the help of a robot (Liquid Handler Quadra 96-320, Tomtec, Hamden, CT). The replica plate was filled with 140 µl of blood-buffer, and after briefly stirring the plates and allowing substrate oxidation in blood-buffer, absorption was measured at 418 nm ($\epsilon_{\text{ABTS}^{+ \cdot}} = 36,000 \text{ M}^{-1} \text{ cm}^{-1}$) in a plate reader (SpectraMax Plus384, Molecular Devices, Sunnyvale, CA). The plates were incubated at room temperature until a green color developed and the absorption was measured again. Relative activities were calculated as the

difference between absorption before and after incubation, and normalized against the parental type used as reference in the corresponding plate (the reference parental types were as follows: 1G, OB-1; 2G, 35H10; 3G, 2E3; 4G, 27C7). To rule out false positives, two consecutive rescreenings were carried out according to our previously described protocol (Maté et al., 2010), but using the blood-buffer assay described above. A third rescreening was introduced to obtain a breakdown of the improvements detected in terms of pH activity profile, Cl^- tolerance, and thermostability.

Laccase Production and Purification

Laccase variants were produced in *S. cerevisiae* (with secretion levels of 8 mg/l) and purified to homogeneity as described previously (Maté et al., 2010). For spectroelectrochemical studies, mutants were overproduced in *Pichia pastoris* after cloning them into the pPICZαA vector under the control of the methanol-inducible AOX1 promoter (*Pichia* expression kit, Life Technologies, Carlsbad, CA). Secretions levels of 43 mg/l were achieved in a 20L autoclavable stainless steel bioreactor (Applikon Biotechnology, Schiedam, The Netherlands).

Measurement of Laccase Activity in Human Plasma and Blood

Human blood was collected in BD Vacutainer blood collection tubes (Plymouth, UK). Blood samples were centrifuged for 10 min at 3,000 rpm to obtain human plasma, discarding the pellet after having carefully extracted the supernatant. Both plasma and blood were supplemented with 10 mM ascorbic acid as substrate and the pH adjusted to 7.4. The activity of the ChU-B mutant in both physiological fluids was determined by measuring oxygen consumption in solution with a Clark electrode. These experiments were performed using the Oxygraph system (Hansatech Instruments, King's Lynn, UK), taking into account the stoichiometry of the reaction: one molecule of O_2 is reduced by oxidizing four substrate molecules. The activity of the ChU-B mutant was further measured in blood-buffer using the same protocol described above and the following substrates: potassium ferrocyanide, $\text{K}_4[\text{Fe}(\text{CN})_6]$ (5 mM final concentration); ABTS (5 mM final concentration); and ascorbic acid (10 mM final concentration). Human plasma and blood manipulation was officially authorized by the Faculty of Health and Society of the Malmö University.

Biochemical Characterization

Determination of Maximum Turnover Rates in Blood-Buffer

The initial turnover rates were determined in blood-buffer (pH 7.4) for ABTS and DMP. A typical reaction started upon the addition of the enzyme (4.7 and 22.7 nM for ABTS and DMP, respectively) to the assay mixture (1.3 mM ABTS or 2.8 mM DMP in blood-buffer). All measurements were performed in triplicate.

pH-Activity Profiles

The pH profiles were determined over a range of pH 2.0–9.0 as described previously (Camarero et al., 2012).

Halide Inhibition

The inhibitory effect of fluoride, chloride and bromide was measured using two laccase substrates (ABTS and DMP) at their corresponding optimal pH activity values (in 100 mM sodium acetate buffer (pH 4.0) for ABTS and 100 mM sodium tartrate buffer (pH 5.0) for DMP), as well as at physiological pH (in 100 mM sodium phosphate buffer, pH 7.4). Inhibition was determined by the I_{50} value (the halide concentration at which only 50% of the initial laccase activity is retained), as the complexity of the plots complicated the extraction of the inhibition constant (K_i). The assay mixture contained 2.4 mM ABTS or DMP, halide (concentrations ranging from 0 to 1100 mM) and purified laccase (0.2 and 1.7 nM for ABTS and DMP, respectively). Each data point represents the mean value determined in at least three independent experiments.

Thermostability

The thermostability of the different laccase samples was estimated by determining their T_{50} values as reported in a previous work (García-Ruiz et al., 2010).

Spectro-Electrochemical Studies

Determination of Laccase Redox Potential

The redox titration of yellow laccase mutants was carried out as described previously for the OB-1 parental type (Mate et al., 2012) with some modifications. Briefly, a microspectroelectrochemical cell containing a gold capillary electrode was used, with the potential controlled by a three-electrode BAS LC-3E potentiostat (Bioanalytical Systems, West Lafayette, IN) where a

$\text{Ag}|\text{AgCl}|\text{KCl}$ electrode was used as reference electrode (BAS) and a platinum wire as counter electrode. The absorbance spectra were monitored on a HR4000 high resolution spectrometer with an effective range of 200–1,100 nm using a DH-2000 light source, both from Ocean Optics (Dunedin, FL). The redox potential of the T1 site of the enzyme was determined by mediated redox titration (MRT) using the spectroelectrochemical set-up described above. A complex mediator system was used for the MRT containing three different redox mediators: $\text{K}_4[\text{Fe}(\text{CN})_6]$ and $\text{K}_4[\text{Mo}(\text{CN})_6]$ with formal redox potentials of +430 mV and +780 mV versus NHE, respectively, and $\text{K}_4[\text{W}(\text{CN})_6]$ with a formal redox potential of +520 mV versus NHE, synthesized as described previously (Leipoldt et al., 1974). All experiments were performed in 50 mM sodium phosphate buffer pH 7.4.

Electrochemical Measurements

Electrochemical experiments were performed with an Autolab PGSTAT30 analyzer controlled by GPES 4.9 software (Eco Chemie, The Netherlands). Experiments were run in a 3-electrode glass cell using a BAS $\text{Ag}|\text{AgCl}|\text{KCl}$ reference electrode (+210 mV versus NHE) and a platinum wire counter electrode. The low-density graphite (LDG electrodes) were abraded with sandpaper and sonicated. A 5 μl drop of laccase (13 mg/ml) was deposited on a clean LDG electrode and after it had been absorbed for 20 min, the electrode was set into the electrochemical cell, which was bubbled with O_2 (g) at 1 atm pressure for 15 min. For electrocatalytic measurements, current densities are expressed relative to the geometric area of the electrodes and potentials are described relative to that of the NHE. Cyclic voltammograms were recorded from +1,100 mV to +210 mV versus NHE using a 10-mV/s scan rate.

Protein Modeling

Protein models of the parental and evolved laccases were generated and analyzed as described previously (Maté et al., 2010).

SUPPLEMENTAL INFORMATION

Supplemental Information includes five tables, three figures, and Supplemental Results and can be found with this article online at <http://dx.doi.org/10.1016/j.chembiol.2013.01.001>.

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